

Form PTO-1390 OFFICE		U.S. DEPARTMENT OF COMMERCE PATENT AND TRADEMARK TRANSMITTAL LETTER TO THE UNITED STATES DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. 371		ATTORNEY'S DOCKET NUMBER GC382-US
INTERNATIONAL APPLICATION NO. PCT/US98/14647		INTERNATIONAL FILING DATE 14 July 1998		U.S. APPLICATION NO. If known, see 37 CFR 1.5) 09/462845 [please provide]
				PRIORITY DATE CLAIMED 15 July 1997
<p>TITLE OF THE INVENTION Proteases from Gram-Positive Organisms</p> <p>APPLICANT(S) FOR DOE/EO/US David A. ESTELL, US Citizen, residing in San Mateo, California 94403, USA</p>				
<p>Applicant herewith submits to the United States Designated/Elected Office (do/eo/us) the following items and other information:</p> <p>1. <input checked="" type="checkbox"/> This is a FIRST submission of items concerning a filing under 35 U.S.C. 371. 2. <input type="checkbox"/> This is a SECOND or SUBSEQUENT submission of items concerning a filing under 35 U.S.C. 371. 3. <input type="checkbox"/> This is an express request to begin national examination procedures (35 U.S.C. 371(f)) at any time rather than delay examination until the expiration of the applicable time limit set in 35 U.S.C. 371(b) and PCT Articles 22 and 39(I). 4. <input checked="" type="checkbox"/> A proper Demand for International Preliminary Examination was made by the 19th month from the earliest claimed priority date. 5. <input checked="" type="checkbox"/> A copy of the International Application as filed (35 U.S.C. 371(c)(2)) a. <input checked="" type="checkbox"/> is transmitted herewith (required only if not transmitted by the International Bureau). b. <input type="checkbox"/> has been transmitted by the International Bureau. c. <input type="checkbox"/> is not required, as the application was filed in the United States Receiving Office (RO/US). 6. <input type="checkbox"/> A translation of the International Application into English (35 U.S.C. 371(c)(2)). 7. <input type="checkbox"/> Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3)) a. <input type="checkbox"/> are transmitted herewith (required only if not transmitted by the International Bureau). b. <input type="checkbox"/> have been transmitted by the International Bureau. c. <input type="checkbox"/> have not been made; however, the time limit for making such amendments has NOT expired. d. <input type="checkbox"/> have not been made and will not be made 8. <input type="checkbox"/> A translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)). 9. <input checked="" type="checkbox"/> An oath or declaration of the inventor/s (35 U.S.C. 371(c)(4)). 10. <input type="checkbox"/> A translation of the annexes to the International Preliminary Examination Report under PCT Article 36 (35 U.S.C. 371(c)(5)).</p> <p>Items 11. to 16 below concern document/s or information included:</p> <p>11. <input type="checkbox"/> An Information Disclosure Statement under 37 CFR 1.97 and 1.98. 12. <input checked="" type="checkbox"/> An assignment document for recording. A separate cover sheet in compliance with 37 CFR 3.28 and 3.31 is included. 13. <input type="checkbox"/> A FIRST preliminary amendment. <input type="checkbox"/> A SECOND or SUBSEQUENT preliminary Amendment 14. <input type="checkbox"/> A substitute specification. 15. <input type="checkbox"/> A change of power of attorney and/or address letter. 16. <input checked="" type="checkbox"/> Other items or information: <ul style="list-style-type: none"> ■ Copy if IPER ■ Copy of Publication of the International Search Report </p>				

U.S. APPLICATION NO. (if known, see 37 CFR 1.5)		INTERNATIONAL APPLICATION NO.	ATTORNEY'S DOCKET NUMBER
[please provide] 097462845		PCT/US98/14647	GC382-US
17. <input checked="" type="checkbox"/> The following fees are submitted:		CALCULATIONS PTO USE ONLY	
BASIC NATIONAL FEE (37 CFR 1.492(2)(1)-(5)):			
Search Report has been prepared by the EPO or JPO		\$ 840	
International preliminary examination fee paid to USPTO (37 CFR 1.482)		\$ 96	
No international preliminary examination fee paid to USPTO (37 CFR 1.482) but international search fee paid to USPTO (37 CFR 1.445(2)(2))		\$	
Neither international preliminary examination fee (37 CFR 1.482) nor international search fee (37 CFR 1.445(2)(2)) paid to USPTO		\$	
International preliminary examination fee paid to USPTO (37 CFR 1.482) and all claims satisfied provisions of PCT Article 33(2)-(4)		\$	
ENTER APPROPRIATE BASIC FEE AMOUNT = \$ 936			
Surcharge of \$130 for furnishing the oath or declaration later than <input type="checkbox"/> 20 <input checked="" type="checkbox"/> 30 months from the earliest claimed priority date (37 CFR 1.492(4)).			
CLAIMS	NUMBER FILED	NUMBER EXTRA	RATE
Total Claims	16	- 20 =	0
		x \$18.00	\$
Independent claims	5	- 3 =	2
		x \$78.00	\$ 156
MULTIPLE DEPENDENT CLAIM(S) (IF APPLICABLE)			
TOTAL OF ABOVE CALCULATIONS = \$1092			
Reduction of 1/2 for filing by small entity, if applicable. Verified Small Entity Statement must also be filed (Note 37 CFR 1.9, 1.27, 1.28)			
SUBTOTAL = \$1092			
Processing fee of \$130 for furnishing the English translation later than <input type="checkbox"/> 20 <input checked="" type="checkbox"/> 30 months from the earliest claimed priority date (37 CFR 1.492(f)).			
TOTAL NATIONAL FEE = \$			
Fee for recording the enclosed assignment (37 CFR 1.21(h)). The assignment must be accompanied by an appropriate cover sheet (37 CFR 3.28, 3.31). \$40.00 per property			
TOTAL FEES ENCLOSED = \$1132			
		Amount to be refunded:	\$
		charged:	\$
<p>a. <input type="checkbox"/> A check in the amount of \$_____ to cover the above fees is enclosed.</p> <p>b. <input checked="" type="checkbox"/> Please charge my Deposit Account No. 07-1048 in the amount of \$1132 to cover the above fees. A duplicate copy of this sheet is enclosed.</p> <p>c. <input checked="" type="checkbox"/> The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 07-1048. A duplicate copy of this sheet is enclosed.</p>			
<p>NOTE: Where an appropriate time limit under 37 CFR 1.494 or 1.495 has not been met, a petition to revive (37 CFR 1.137(a) or (b)) must be filed and granted to restore the application to pending status.</p>			
<p>SEND ALL CORRESPONDENCE TO:</p> <p>Jeffery D. FRAZIER GENENCOR INTERNATIONAL, INC. 925 PAGE MILL ROAD PALO ALTO, CALIFORNIA 94304-1013</p> <p><i>Jeffery D. Frazier</i> SIGNATURE</p> <p>Jeffery D. FRAZIER <small>(printed name)</small></p>			
<p>Tel. 650-846-7655 7615 Fax 650-845-6504</p> <p>Registration Number: 34,601</p>			

PROTEASES FROM GRAM-POSITIVE ORGANISMSFIELD OF THE INVENTION

The present invention relates to serine proteases derived from gram-positive microorganisms. The present invention provides nucleic acid and amino acid sequences of serine protease 1, 2, 3, 4 and 5 identified in *Bacillus*. The present invention also provides methods for the production of serine protease 1, 2, 3, 4 and 5 in host cells as well as the production of heterologous proteins in a host cell having a mutation or deletion of part or all of at least one of the serine proteases of the present invention.

BACKGROUND OF THE INVENTION

Gram-positive microorganisms, such as members of the group *Bacillus*, have been used for large-scale industrial fermentation due, in part, to their ability to secrete their fermentation products into the culture media. In gram-positive bacteria, secreted proteins are exported across a cell membrane and a cell wall, and then are subsequently released into the external media usually maintaining their native conformation.

Various gram-positive microorganisms are known to secrete extracellular and/or intracellular protease at some stage in their life cycles. Many proteases are produced in large quantities for industrial purposes. A negative aspect of the presence of proteases in gram-positive organisms is their contribution to the overall degradation of secreted heterologous or foreign proteins.

The classification of proteases found in microorganisms is based on their catalytic mechanism which results in four groups: the serine proteases; metalloproteases; cysteine proteases; and aspartic proteases. These categories can be distinguished by their sensitivity to various inhibitors. For example, the serine proteases are inhibited by phenylmethylsulfonylfluoride (PMSF) and diisopropylfluorophosphate (DIFP); the metalloproteases by chelating agents; the cysteine enzymes by iodoacetamide and heavy metals and the aspartic proteases by pepstatin. The serine proteases have alkaline pH optima, the metalloproteases are optimally active around neutrality, and the cysteine and aspartic enzymes have acidic pH optima (*Biotechnology Handbooks, Bacillus*, vol. 2, edited by Harwood, 1989 Plenum Press, New York).

Proteolytic enzymes that are dependent upon a serine residue for catalytic activity are called serine proteases. As described in *Methods in Enzymology*, vol. 244, Academic Press, Inc. 1994, page 21, serine proteases of the family S9 have the catalytic residue triad "Ser-Asp-His with conservation of amino acids around them.

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SUMMARY OF THE INVENTION

The present invention relates to the unexpected discovery of five heretofore unknown or unrecognized S9 type serine proteases found in uncharacterized translated genomic nucleic acid sequences of *Bacillus subtilis*, designated herein as SP1, SP2, SP3, SP4 and SP5 having the nucleic acid and amino acid as shown in the Figures. The present invention is based, in part, upon the presence the amino acid triad S-D-H in the five serine proteases, as well as amino acid conservation around the triad. The present invention is also based in part upon the heretofore uncharacterized or unrecognized overall amino acid relatedness that SP1, SP2, SP3, SP4 and SP5 have with the serine protease dipeptidyl-amino peptidase B from yeast (DAP) and with each other.

The present invention provides isolated polynucleotide and amino acid sequences for SP1, SP2, SP3, SP4 and SP5. Due to the degeneracy of the genetic code, the present invention encompasses any nucleic acid sequence that encodes the SP1, SP2, SP3, SP4 and SP5 deduced amino acid sequences shown in Figures 2A-2B-Figure 6, respectively.

The present invention encompasses amino acid variations of *B. subtilis* SP1, SP2, SP3, SP4 and SP5 disclosed herein that have proteolytic activity. *B. subtilis* SP1, SP2, SP3, SP4 and SP5, as well as proteolytically active amino acid variations thereof, have application in cleaning compositions. In one aspect of the present invention, SP1, SP2, SP3, SP4 and SP5 obtainable from a gram-positive microorganism are produced on an industrial fermentation scale in a microbial host expression system. In another aspect, isolated and purified SP1, SP2, SP3, SP4 or SP5 obtainable from a gram-positive microorganism is used in compositions of matter intended for cleaning purposes, such as detergents. Accordingly, the present invention provides a cleaning composition comprising at least one of SP1, SP2, SP3, SP4 and SP5 obtainable from a gram-positive microorganism. The serine protease may be used alone in the cleaning composition or in combination with other enzymes and/or mediators or enhancers.

The production of desired heterologous proteins or polypeptides in gram-positive microorganisms may be hindered by the presence of one or more proteases which degrade the produced heterologous protein or polypeptide. Therefore, the present invention also encompasses gram-positive microorganism having a mutation or deletion of part or all of the gene encoding SP1, SP2, SP3, SP4 and/or SP5, which results in the inactivation of their proteolytic activity, either alone or in combination with deletions or mutations in other proteases, such as apr, npr, epr, mpr for example, or other proteases known to those of skill in the art. In one embodiment of the present invention, the gram-positive organism is a member of the genus *Bacillus*. In another embodiment, the *Bacillus* is *Bacillus subtilis*.

In another aspect, the gram-positive microorganism host having one or more deletions or mutations in a serine protease of the present invention is further genetically

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engineered to produce a desired protein. In one embodiment of the present invention, the desired protein is heterologous to the gram-positive host cell. In another embodiment, the desired protein is homologous to the host cell. The present invention encompasses a gram-positive host cell having a deletion or interruption of the naturally occurring nucleic acid encoding the homologous protein, such as a protease, and having nucleic acid encoding the homologous protein or a variant thereof re-introduced in a recombinant form. In another embodiment, the host cell produces the homologous protein. Accordingly, the present invention also provides methods and expression systems for reducing degradation of heterologous or homologous proteins produced in gram-positive microorganisms comprising the steps of obtaining a *Bacillus* host cell comprising nucleic acid encoding said heterologous protein wherein said host cell contains a mutation or deletion in at least one of the genes encoding SP1, SP2, SP3, SP4 and SP5; and growing said *Bacillus* host cell under conditions suitable for the expression of said heterologous protein. The gram-positive microorganism may be normally sporulating or non-sporulating.

The present invention provides methods for detecting gram positive microorganism homologs of *B. subtilis* SP1, SP2, SP3, SP4 and SP5 that comprises hybridizing part or all of the nucleic acid encoding *B. subtilis* SP1, SP2, SP3, SP4 and SP5 with nucleic acid derived from gram-positive organisms, either of genomic or cDNA origin.

BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1A-1C shows the DNA (SEQ ID NO:1) and deduced amino acid sequence (SEQ ID NO:2) for SP1 (YUXL).

Figures 2A-2B show an amino acid alignment between DAP (dap2_yeast) (SEQ ID NO:3) and SP1 (YUXL). For Figures 2A-2B, 3 and 4, the amino acid triad S-D-H is indicated.

Figure 3 shows an amino acid alignment between SP1 (YUXL) (SEQ ID NO:2) and SP2 (YTMA) (SEQ ID NO:5).

Figure 4 shows an amino acid alignment between SP1 (YUXL) (SEQ ID NO:2) and SP3 (YITV) (SEQ ID NO:7).

Figure 5 shows an amino acid alignment between SP1 (YUXL) (SEQ ID NO:2) and SP4 (YQKD) (SEQ ID NO:9).

Figure 6 shows an amino acid alignment between SP1 (YUXL) (SEQ ID NO:2) and SP5 (CAH) (SEQ ID NO:10).

Figures 7A-7B shows the DNA (SEQ ID NO:4) and deduced amino acid sequence for SP2 (YTMA) (SEQ ID NO:5).

Figures 8A-8B shows the DNA (SEQ ID NO:6) and deduced amino acid sequence for SP3 (YITV) (SEQ ID NO:7).

Figures 9A-9B shows the DNA (SEQ ID NO:8) and deduced amino acid sequence for SP4 (YQKD) (SEQ ID NO:9).

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

Definitions — As used herein, the genus *Bacillus* includes all members known to those of skill in the art, including but not limited to *B. subtilis*, *B. licheniformis*, *B. lentus*, *B. brevis*, *B. stearothermophilus*, *B. alkalophilus*, *B. amyloliquefaciens*, *B. coagulans*, *B. cibulans*, *B. lautus* and *B. thuringiensis*.

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The present invention encompasses novel SP1, SP2, SP3, SP4 and SP5 from gram positive organisms. In a preferred embodiment, the gram-positive organisms is a *Bacillus*. In another preferred embodiment, the gram-positive organism is *Bacillus subtilis*. As used herein, *B. subtilis* SP1 (YuxL) refers to the DNA and deduced amino acid sequence shown in Figures 1A-1C and Figures 2A-2B; SP2 (YtmA) refers to the DNA and deduced amino acid sequence shown in Figures 7A-7B and Figure 3; SP3 (YitV) refers to the DNA and deduced amino acid sequence shown in Figures 8A-8B and Figure 4; SP4 (YqkD) refers to the DNA and deduced amino acid sequence shown in Figures 9A-9B and Figure 5; and SP5 (CAH) refers to the deduced amino acid sequence shown in Figure 6. The present invention encompasses amino acid variations of the *B. subtilis* amino acid sequences of SP1, SP2, SP3, SP4 and SP5 that have proteolytic activity. Such proteolytic amino acid variants can be used in cleaning compositions.

As used herein, "nucleic acid" refers to a nucleotide or polynucleotide sequence, and fragments or portions thereof, and to DNA or RNA of genomic or synthetic origin which may be double-stranded or single-stranded, whether representing the sense or antisense strand. As used herein "amino acid" refers to peptide or protein sequences or portions thereof. A "polynucleotide homolog" as used herein refers to a novel gram-positive microorganism polynucleotide that has at least 80%, at least 90% and at least 95% identity to *B. subtilis* SP1, SP2, SP3, SP4 or SP5, or which is capable of hybridizing to *B. subtilis* SP1, SP2, SP3, SP4 or SP5 under conditions of high stringency and which encodes an amino acid sequence having serine protease activity.

The terms "isolated" or "purified" as used herein refer to a nucleic acid or amino acid that is removed from at least one component with which it is naturally associated.

As used herein, the term "heterologous protein" refers to a protein or polypeptide that does not naturally occur in a gram-positive host cell. Examples of heterologous proteins include enzymes such as hydrolases including proteases, cellulases, amylases, carbohydrases, and lipases; isomerases such as racemases, epimerases, tautomerases, or mutases; transferases, kinases and phosphatases. The heterologous gene may encode therapeutically significant proteins or peptides, such as growth factors, cytokines, ligands, receptors and inhibitors, as well as vaccines and antibodies. The gene may encode commercially important industrial proteins or peptides, such as proteases, carbohydrases such as amylases and glucoamylases, cellulases, oxidases and lipases. The gene of interest may be a naturally occurring gene, a mutated gene or a synthetic gene.

The term "homologous protein" refers to a protein or polypeptide native or naturally occurring in a gram-positive host cell. The invention includes host cells producing the homologous protein via recombinant DNA technology. The present invention encompasses

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a gram-positive host cell having a deletion or interruption of the nucleic acid encoding the naturally occurring homologous protein, such as a protease, and having nucleic acid encoding the homologous protein, or a variant thereof re-introduced in a recombinant form. In another embodiment, the host cell produces the homologous protein.

As used herein, the term "overexpressing" when referring to the production of a protein in a host cell means that the protein is produced in greater amounts than its production in its naturally occurring environment.

As used herein, the phrase "proteolytic activity" refers to a protein that is able to hydrolyze a peptide bond. Enzymes having proteolytic activity are described in Enzyme Nomenclature, 1992, edited Webb Academic Press, Inc.

Detailed Description of the Preferred Embodiments

The unexpected discovery of the serine proteases SP1, SP2, SP3, SP4 and SP5 in *B. subtilis* provides a basis for producing host cells, expression methods and systems which can be used to prevent the degradation of recombinantly produced heterologous proteins. In a preferred embodiment, the host cell is a gram-positive host cell that has a deletion or mutation in the naturally occurring serine protease said mutation resulting in the complete deletion or inactivation of the production by the host cell of the proteolytic serine protease gene product. In another embodiment of the present invention, the host cell is additionally genetically engineered to produce a desired protein or polypeptide.

It may also be desired to genetically engineer host cells of any type to produce a gram-positive serine protease SP1, SP2, SP3, SP4 or SP5. Such host cells are used in large scale fermentation to produce large quantities of the serine protease which may be isolated or purified and used in cleaning products, such as detergents.

I. Serine Protease Nucleic Acid and Amino Acid Sequences

The SP1, SP2, SP3 and SP4 polynucleotides having the sequences as shown in the Figures encode the *Bacillus subtilis* serine SP1, SP2, SP3, and SP4. As will be understood by the skilled artisan, due to the degeneracy of the genetic code, a variety of polynucleotides can encode the *Bacillus* SP1, SP2, SP3, SP4 and SP5. The present invention encompasses all such polynucleotides.

The present invention encompasses novel SP1, SP2, SP3, SP4 and SP5 polynucleotide homologs encoding gram-positive microorganism serine proteases SP1, SP2, SP3, SP4 and SP5, respectively, which have at least 80%, or at least 90% or at least 95% identity to *B. subtilis* as long as the homolog encodes a protein that has proteolytic activity.

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Gram-positive polynucleotide homologs of *B. subtilis* SP1, SP2, SP3, SP4 or SP5 may be obtained by standard procedures known in the art from, for example, cloned DNA (e.g., a DNA "library"), genomic DNA libraries, by chemical synthesis once identified, by cDNA cloning, or by the cloning of genomic DNA, or fragments thereof, purified from a desired cell. (See, for example, Sambrook et al., 1989, Molecular Cloning, A Laboratory Manual, 2d Ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, New York; Glover, D.M. (ed.), 1985, DNA Cloning: A Practical Approach, MRL Press, Ltd., Oxford, U.K. Vol. I, II.) A preferred source is from genomic DNA. Nucleic acid sequences derived from genomic DNA may contain regulatory regions in addition to coding regions. Whatever the source, the isolated serine protease gene should be molecularly cloned into a suitable vector for propagation of the gene.

In the molecular cloning of the gene from genomic DNA, DNA fragments are generated, some of which will encode the desired gene. The DNA may be cleaved at specific sites using various restriction enzymes. Alternatively, one may use DNase in the presence of manganese to fragment the DNA, or the DNA can be physically sheared, as for example, by sonication. The linear DNA fragments can then be separated according to size by standard techniques, including but not limited to, agarose and polyacrylamide gel electrophoresis and column chromatography.

Once the DNA fragments are generated, identification of the specific DNA fragment containing the SP1, SP2, SP3, SP4 or SP5 may be accomplished in a number of ways. For example, a *B. subtilis* SP1, SP2, SP3, SP4 or SP5 gene of the present invention or its specific RNA, or a fragment thereof, such as a probe or primer, may be isolated and labeled and then used in hybridization assays to detect a gram-positive SP1, SP2, SP3, SP4 or SP5 gene. (Benton, W. and Davis, R., 1977, *Science* 196:180; Grunstein, M. And Hogness, D., 1975, *Proc. Natl. Acad. Sci. USA* 72:3961). Those DNA fragments sharing substantial sequence similarity to the probe will hybridize under stringent conditions.

Accordingly, the present invention provides a method for the detection of gram-positive SP1, SP2, SP3, SP4 or SP5 polynucleotide homologs which comprises hybridizing part or all of a nucleic acid sequence of *B. subtilis* SP1, SP2, SP3, SP4 or SP5 with gram-positive microorganism nucleic acid of either genomic or cDNA origin.

Also included within the scope of the present invention are gram-positive microorganism polynucleotide sequences that are capable of hybridizing to the nucleotide sequence of *B. subtilis* SP1, SP2, SP3, SP4 or SP5 under conditions of intermediate to maximal stringency. Hybridization conditions are based on the melting temperature (Tm) of the nucleic acid binding complex, as taught in Berger and Kimmel (1987, *Guide to Molecular Cloning Techniques*, Methods in Enzymology, Vol 152, Academic Press, San Diego CA) incorporated herein by reference, and confer a defined "stringency" as explained below.

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"Maximum stringency" typically occurs at about $T_m - 5^\circ\text{C}$ (5°C below the T_m of the probe); "high stringency" at about 5°C to 10°C below T_m ; "intermediate stringency" at about 10°C to 20°C below T_m ; and "low stringency" at about 20°C to 25°C below T_m . As will be understood by those of skill in the art, a maximum stringency hybridization can be used to identify or detect identical polynucleotide sequences while an intermediate or low stringency hybridization can be used to identify or detect polynucleotide sequence homologs.

The term "hybridization" as used herein shall include "the process by which a strand of nucleic acid joins with a complementary strand through base pairing" (Coombs J (1994) Dictionary of Biotechnology, Stockton Press, New York NY).

10 The process of amplification as carried out in polymerase chain reaction (PCR) technologies is described in Dieffenbach CW and GS Dveksler (1995, PCR Primer, a Laboratory Manual, Cold Spring Harbor Press, Plainview NY). A nucleic acid sequence of at least about 10 nucleotides and as many as about 60 nucleotides from *B. subtilis* SP1, SP2, SP3, SP4 or SP5 preferably about 12 to 30 nucleotides, and more preferably about 15 20-25 nucleotides can be used as a probe or PCR primer.

15 The *B. subtilis* amino acid sequences SP1, SP2, SP3, SP4 and SP5 (shown in Figures 2A-2B through Figure 6) were identified via a FASTA search of *Bacillus subtilis* genomic nucleic acid sequences. *B. subtilis* SP1 (YuxL) was identified by its structural homology to the serine protease DAP classified as an S9 type serine protease, designated in Figures 2A-2B as "dap2_yeast". As shown in Figures 2A-2B, SP1 has the amino acid dyad "S-D-H" indicated. Conservation of amino acids around each residue is noted in Figures 2A-2B through Figure 6. SP2 (YtmA); SP3 (YitV); SP4 (YqkD) and SP5 (CAH) were identified upon by their structural and overall amino acid homology to SP1 (YuxL). SP1 and SP4 were described in Parsot and Kebayashi, respectively, but were not characterized as 20 serine proteases or serine proteases of the S9 family.

II. Expression Systems

25 The present invention provides host cells, expression methods and systems for the enhanced production and secretion of desired heterologous or homologous proteins in gram-positive microorganisms. In one embodiment, a host cell is genetically engineered to have a deletion or mutation in the gene encoding a gram-positive SP1, SP2, SP3, SP4 or SP5 such that the respective activity is deleted. In an alternative embodiment of the present invention, a gram-positive microorganism is genetically engineered to produce a 30 serine protease of the present invention.

Inactivation of a gram-positive serine protease in a host cell

Producing an expression host cell incapable of producing the naturally occurring serine protease necessitates the replacement and/or inactivation of the naturally occurring gene from the genome of the host cell. In a preferred embodiment, the mutation is a non-reverting mutation.

One method for mutating nucleic acid encoding a gram-positive serine protease is to clone the nucleic acid or part thereof, modify the nucleic acid by site directed mutagenesis and reintroduce the mutated nucleic acid into the cell on a plasmid. By homologous recombination, the mutated gene may be introduced into the chromosome. In the parent host cell, the result is that the naturally occurring nucleic acid and the mutated nucleic acid are located in tandem on the chromosome. After a second recombination, the modified sequence is left in the chromosome having thereby effectively introduced the mutation into the chromosomal gene for progeny of the parent host cell.

Another method for inactivating the serine protease proteolytic activity is through deleting the chromosomal gene copy. In a preferred embodiment, the entire gene is deleted, the deletion occurring in such a way as to make reversion impossible. In another preferred embodiment, a partial deletion is produced, provided that the nucleic acid sequence left in the chromosome is too short for homologous recombination with a plasmid encoded serine protease gene. In another preferred embodiment, nucleic acid encoding the catalytic amino acid residues are deleted.

Deletion of the naturally occurring gram-positive microorganism serine protease can be carried out as follows. A serine protease gene including its 5' and 3' regions is isolated and inserted into a cloning vector. The coding region of the serine protease gene is deleted from the vector *in vitro*, leaving behind a sufficient amount of the 5' and 3' flanking sequences to provide for homologous recombination with the naturally occurring gene in the parent host cell. The vector is then transformed into the gram-positive host cell. The vector integrates into the chromosome via homologous recombination in the flanking regions. This method leads to a gram-positive strain in which the protease gene has been deleted.

The vector used in an integration method is preferably a plasmid. A selectable marker may be included to allow for ease of identification of desired recombinant microorganisms. Additionally, as will be appreciated by one of skill in the art, the vector is preferably one which can be selectively integrated into the chromosome. This can be achieved by introducing an inducible origin of replication, for example, a temperature sensitive origin into the plasmid. By growing the transformants at a temperature to which the origin of replication is sensitive, the replication function of the plasmid is inactivated, thereby providing a means for selection of chromosomal integrants. Integrants may be

selected for growth at high temperatures in the presence of the selectable marker, such as an antibiotic. Integration mechanisms are described in WO 88/06623.

Integration by the Campbell-type mechanism can take place in the 5' flanking region of the protease gene, resulting in a protease positive strain carrying the entire plasmid vector in the chromosome in the serine protease locus. Since illegitimate recombination will give different results it will be necessary to determine whether the complete gene has been deleted, such as through nucleic acid sequencing or restriction maps.

Another method of inactivating the naturally occurring serine protease gene is to mutagenize the chromosomal gene copy by transforming a gram-positive microorganism with oligonucleotides which are mutagenic. Alternatively, the chromosomal serine protease gene can be replaced with a mutant gene by homologous recombination.

The present invention encompasses host cells having additional protease deletions or mutations, such as deletions or mutations in apr, npr, epr, mpr and others known to those of skill in the art. United States Patent 5,264,366 discloses *Bacillus* host cells having a deletion of apr and npr; United States Patent 5,585,253 discloses *Bacillus* host cells having a deletion of epr; Margot et al., 1996, *Microbiology* 142: 3437-3444 disclose host cells having a deletion in wpr and EP patent 0369817 discloses *Bacillus* host cells having a deletion of mpr.

III. Production of Serine protease

For production of serine protease in a host cell, an expression vector comprising at least one copy of nucleic acid encoding a gram-positive microorganism SP1, SP2, SP3, SP4 or SP5, and preferably comprising multiple copies, is transformed into the host cell under conditions suitable for expression of the serine protease. In accordance with the present invention, polynucleotides which encode a gram-positive microorganism SP1, SP2, SP3, SP4 or SP5, or fragments thereof, or fusion proteins or polynucleotide homolog sequences that encode amino acid variants of B. SP1, SP2, SP3, SP4 or SP5, may be used to generate recombinant DNA molecules that direct their expression in host cells. In a preferred embodiment, the gram-positive host cell belongs to the genus *Bacillus*. In another preferred embodiment, the gram positive host cell is *B. subtilis*.

As will be understood by those of skill in the art, it may be advantageous to produce polynucleotide sequences possessing non-naturally occurring codons. Codons preferred by a particular gram-positive host cell (Murray E et al (1989) *Nuc Acids Res* 17:477-508) can be selected, for example, to increase the rate of expression or to produce recombinant RNA transcripts having desirable properties, such as a longer half-life, than transcripts produced from naturally occurring sequence.

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Altered SP1, SP2, SP3, SP4 or SP5 polynucleotide sequences which may be used in accordance with the invention include deletions, insertions or substitutions of different nucleotide residues resulting in a polynucleotide that encodes the same or a functionally equivalent SP1, SP2, SP3, SP4 or SP5 homolog, respectively. As used herein a "deletion" is defined as a change in either nucleotide or amino acid sequence in which one or more nucleotides or amino acid residues, respectively, are absent.

As used herein an "insertion" or "addition" is that change in a nucleotide or amino acid sequence which has resulted in the addition of one or more nucleotides or amino acid residues, respectively, as compared to the naturally occurring SP1, SP2, SP3, SP4 or SP5.

As used herein "substitution" results from the replacement of one or more nucleotides or amino acids by different nucleotides or amino acids, respectively.

The encoded protein may also show deletions, insertions or substitutions of amino acid residues which produce a silent change and result in a functionally SP1, SP2, SP3, SP4 or SP5 variant. Deliberate amino acid substitutions may be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity, and/or the amphipathic nature of the residues as long as the variant retains the ability to modulate secretion. For example, negatively charged amino acids include aspartic acid and glutamic acid; positively charged amino acids include lysine and arginine; and amino acids with uncharged polar head groups having similar hydrophilicity values include leucine, isoleucine, valine; glycine, alanine; asparagine, glutamine; serine, threonine, phenylalanine, and tyrosine.

The SP1, SP2, SP3, SP4 or SP5 polynucleotides of the present invention may be engineered in order to modify the cloning, processing and/or expression of the gene product. For example, mutations may be introduced using techniques which are well known in the art, eg, site-directed mutagenesis to insert new restriction sites, to alter glycosylation patterns or to change codon preference, for example.

In one embodiment of the present invention, a gram-positive microorganism SP1, SP2, SP3, SP4 or SP5 polynucleotide may be ligated to a heterologous sequence to encode a fusion protein. A fusion protein may also be engineered to contain a cleavage site located between the serine protease nucleotide sequence and the heterologous protein sequence, so that the serine protease may be cleaved and purified away from the heterologous moiety.

IV. Vector Sequences

Expression vectors used in expressing the serine proteases of the present invention in gram-positive microorganisms comprise at least one promoter associated with a serine protease selected from the group consisting of SP1, SP2, SP3, SP4 and SP5, which

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promoter is functional in the host cell. In one embodiment of the present invention, the promoter is the wild-type promoter for the selected serine protease and in another embodiment of the present invention, the promoter is heterologous to the serine protease, but still functional in the host cell. In one preferred embodiment of the present invention, the nucleic acid encoding the serine protease is stably integrated into the microorganism genome.

In a preferred embodiment, the expression vector contains a multiple cloning site cassette which preferably comprises at least one restriction endonuclease site unique to the vector, to facilitate ease of nucleic acid manipulation. In a preferred embodiment, the vector also comprises one or more selectable markers. As used herein, the term selectable marker refers to a gene capable of expression in the gram-positive host which allows for ease of selection of those hosts containing the vector. Examples of such selectable markers include but are not limited to antibiotics, such as, erythromycin, actinomycin, chloramphenicol and tetracycline.

V. Transformation

A variety of host cells can be used for the production of SP1, SP2, SP3, SP4 or SP5 including bacterial, fungal, mammalian and insects cells. General transformation procedures are taught in Current Protocols In Molecular Biology (vol. 1, edited by Ausubel et al., John Wiley & Sons, Inc. 1987, Chapter 9) and include calcium phosphate methods, transformation using DEAE-Dextran and electroporation. Plant transformation methods are taught in Rodriguez (WO 95/14099, published 26 May 1995).

In a preferred embodiment, the host cell is a gram-positive microorganism and in another preferred embodiment, the host cell is *Bacillus*. In one embodiment of the present invention, nucleic acid encoding one or more serine protease(s) of the present invention is introduced into a host cell via an expression vector capable of replicating within the host cell. Suitable replicating plasmids for *Bacillus* are described in Molecular Biological Methods for *Bacillus*, Ed. Harwood and Cutting, John Wiley & Sons, 1990, hereby expressly incorporated by reference; see chapter 3 on plasmids. Suitable replicating plasmids for *B. subtilis* are listed on page 92.

In another embodiment, nucleic acid encoding a serine protease(s) of the present invention is stably integrated into the microorganism genome. Preferred host cells are gram-positive host cells. Another preferred host is *Bacillus*. Another preferred host is *Bacillus subtilis*. Several strategies have been described in the literature for the direct cloning of DNA in *Bacillus*. Plasmid marker rescue transformation involves the uptake of a donor plasmid by competent cells carrying a partially homologous resident plasmid

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(Contente *et al.*, *Plasmid* 2:555-571 (1979); Haima *et al.*, *Mol. Gen. Genet.* 223:185-191 (1990); Weinrauch *et al.*, *J. Bacteriol.* 154(3):1077-1087 (1983); and Weinrauch *et al.*, *J. Bacteriol.* 169(3):1205-1211 (1987)). The incoming donor plasmid recombines with the homologous region of the resident "helper" plasmid in a process that mimics chromosomal transformation.

Transformation by protoplast transformation is described for *B. subtilis* in Chang and Cohen, (1979) *Mol. Gen. Genet.* 168:111-115; for *B. megaterium* in Vorobjeva *et al.*, (1980) *FEMS Microbiol. Letters* 7:261-263; for *B. amyloliquefaciens* in Smith *et al.*, (1986) *Appl. and Env. Microbiol.* 51:634; for *B. thuringiensis* in Fisher *et al.*, (1981) *Arch. Microbiol.* 139:213-217; for *B. sphaericus* in McDonald (1984) *J. Gen. Microbiol.* 130:203; and *B. larvae* in Bakhiet *et al.*, (1985) 49:577. Mann *et al.*, (1986, *Current Microbiol.* 13:131-135) report on transformation of *Bacillus* protoplasts and Holubova, (1985) *Folia Microbiol.* 30:97 disclose methods for introducing DNA into protoplasts using DNA containing liposomes.

VI. Identification of Transformants

Whether a host cell has been transformed with a mutated or a naturally occurring gene encoding a gram-positive SP1, SP2, SP3, SP4 or SP5, detection of the presence/absence of marker gene expression can suggest whether the gene of interest is present. However, its expression should be confirmed. For example, if the nucleic acid encoding a serine protease is inserted within a marker gene sequence, recombinant cells containing the insert can be identified by the absence of marker gene function. Alternatively, a marker gene can be placed in tandem with nucleic acid encoding the serine protease under the control of a single promoter. Expression of the marker gene in response to induction or selection usually indicates expression of the serine protease as well.

Alternatively, host cells which contain the coding sequence for a serine protease and express the protein may be identified by a variety of procedures known to those of skill in the art. These procedures include, but are not limited to, DNA-DNA or DNA-RNA hybridization and protein bioassay or immunoassay techniques which include membrane-based, solution-based, or chip-based technologies for the detection and/or quantification of the nucleic acid or protein.

The presence of the cysteine polynucleotide sequence can be detected by DNA-DNA or DNA-RNA hybridization or amplification using probes, portions or fragments of *B. subtilis* SP1, SP2, SP3, SP4 or SP5.

VII. Assay of Protease Activity

There are various assays known to those of skill in the art for detecting and measuring protease activity. There are assays based upon the release of acid-soluble peptides from casein or hemoglobin measured as absorbance at 280 nm or colorimetrically using the Folin method (Bergmeyer, et al., 1984, *Methods of Enzymatic Analysis* vol. 5, *Peptidases, Proteinases and their Inhibitors*, Verlag Chemie, Weinheim). Other assays involve the solubilization of chromogenic substrates (Ward, 1983, *Proteinases, in Microbial Enzymes and Biotechnology* (W.M. Fogarty, ed.), Applied Science, London, pp. 251-317).

VIII. Secretion of Recombinant Proteins

Means for determining the levels of secretion of a heterologous or homologous protein in a gram-positive host cell and detecting secreted proteins include, using either polyclonal or monoclonal antibodies specific for the protein. Examples include enzyme-linked immunosorbent assay (ELISA), radioimmunoassay (RIA) and fluorescent activated cell sorting (FACS). These and other assays are described, among other places, in Hampton R et al (1990, *Serological Methods, a Laboratory Manual*, APS Press, St Paul MN) and Maddox DE et al (1983, *J Exp Med* 158:1211).

A wide variety of labels and conjugation techniques are known by those skilled in the art and can be used in various nucleic and amino acid assays. Means for producing labeled hybridization or PCR probes for detecting specific polynucleotide sequences include oligolabeling, nick translation, end-labeling or PCR amplification using a labeled nucleotide. Alternatively, the nucleotide sequence, or any portion of it, may be cloned into a vector for the production of an mRNA probe. Such vectors are known in the art, are commercially available, and may be used to synthesize RNA probes *in vitro* by addition of an appropriate RNA polymerase such as T7, T3 or SP6 and labeled nucleotides.

A number of companies such as Pharmacia Biotech (Piscataway NJ), Promega (Madison WI), and US Biochemical Corp (Cleveland OH) supply commercial kits and protocols for these procedures. Suitable reporter molecules or labels include those radionuclides, enzymes, fluorescent, chemiluminescent, or chromogenic agents as well as substrates, cofactors, inhibitors, magnetic particles and the like. Patents teaching the use of such labels include US Patents 3,817,837; 3,850,752; 3,939,350; 3,996,345; 4,277,437; 4,275,149 and 4,366,241. Also, recombinant immunoglobulins may be produced as shown in US Patent No. 4,816,567 and incorporated herein by reference.

IX. Purification of Proteins

Gram positive host cells transformed with polynucleotide sequences encoding heterologous or homologous protein may be cultured under conditions suitable for the expression and recovery of the encoded protein from cell culture. The protein produced by

a recombinant gram-positive host cell comprising a serine protease of the present invention will be secreted into the culture media. Other recombinant constructions may join the heterologous or homologous polynucleotide sequences to nucleotide sequence encoding a polypeptide domain which will facilitate purification of soluble proteins (Kroll DJ et al (1993) DNA Cell Biol 12:441-53).

Such purification facilitating domains include, but are not limited to, metal chelating peptides such as histidine-tryptophan modules that allow purification on immobilized metals (Porath J (1992) Protein Expr Purif 3:263-281), protein A domains that allow purification on immobilized immunoglobulin, and the domain utilized in the FLAGS extension/affinity purification system (Immunex Corp, Seattle WA). The inclusion of a cleavable linker sequence such as Factor XA or enterokinase (Invitrogen, San Diego CA) between the purification domain and the heterologous protein can be used to facilitate purification.

X. Uses of The Present Invention

Genetically Engineered Host Cells

The present invention provides genetically engineered host cells comprising preferably non-revertable mutations or deletions in the naturally occurring gene encoding one or more of SP1, SP2, SP3, SP4 or SP5 such that the proteolytic activity is diminished or deleted altogether. The host cell may contain additional protease deletions, such as 20 deletions of the mature subtilisin protease and/or mature neutral protease disclosed in United States Patent No. 5,264,366.

In a preferred embodiment, the host cell is genetically engineered to produce a desired protein or polypeptide. In a preferred embodiment the host cell is a *Bacillus*. In another preferred embodiment, the host cell is a *Bacillus subtilis*.

25 In an alternative embodiment, a host cell is genetically engineered to produce a gram-positive SP1, SP2, SP3, SP4 or SP5. In a preferred embodiment, the host cell is grown under large scale fermentation conditions, the SP1, SP2, SP3, SP4 or SP5 is isolated and/or purified and used in cleaning compositions such as detergents. WO 95/10615 discloses detergent formulation. A serine protease of the present invention can be 30 useful in formulating various cleaning compositions. A number of known compounds are suitable surfactants useful in compositions comprising the serine protease of the invention. These include nonionic, anionic, cationic, anionic or zwitterionic detergents, as disclosed in US 4,404,128 and US 4,261,868. A suitable detergent formulation is that described in Example 7 of US Patent 5,204,015. The art is familiar with the different formulations which 35 can be used as cleaning compositions. In addition, a serine protease of the present invention can be used, for example, in bar or liquid soap applications, dishcare formulations,

contact lens cleaning solutions or products, peptide hydrolysis, waste treatment, textile applications, as fusion-cleavage enzymes in protein production, etc. A serine protease of the present invention may provide enhanced performance in a detergent composition (as compared to another detergent protease). As used herein, enhanced performance in a detergent is defined as increasing cleaning of certain enzyme sensitive stains such as grass or blood, as determined by usual evaluation after a standard wash cycle.

A serine protease of the present invention can be formulated into known powdered and liquid detergents having pH between 6.5 and 12.0 at levels of about .01 to about 5% (preferably .1% to .5%) by weight. These detergent cleaning compositions can also include other enzymes such as known proteases, amylases, cellulases, lipases or endoglycosidases, as well as builders and stabilizers.

The addition of a serine protease to conventional cleaning compositions does not create any special use limitation. In other words, any temperature and pH suitable for the detergent is also suitable for the present compositions as long as the pH is within the above range, and the temperature is below the described serine protease denaturing temperature. In addition, a serine protease of the present invention can be used in a cleaning composition without detergents, again either alone or in combination with builders and stabilizers.

One aspect of the invention is a composition for the treatment of a textile that includes a serine protease of the present invention. The composition can be used to treat for example silk or wool as described in publications such as RD 216,034; EP 134,267; US 4,533,359; and EP 344,259.

Proteases can be included in animal feed such as part of animal feed additives as described in, for example, US 5,612,055; US 5,314,692; and US 5,147,642.

Polynucleotides

A *B.subtilis* SP1, SP2, SP3, SP4 or SP5 polynucleotide, or any part thereof, provides the basis for detecting the presence of gram-positive microorganism polynucleotide homologs through hybridization techniques and PCR technology.

Accordingly, one aspect of the present invention is to provide for nucleic acid hybridization and PCR probes which can be used to detect polynucleotide sequences, including genomic and cDNA sequences, encoding gram-positive SP1, SP2, SP3, SP4 or SP5 or portions thereof.

The manner and method of carrying out the present invention may be more fully understood by those of skill in the art by reference to the following examples, which

examples are not intended in any manner to limit the scope of the present invention or of the claims directed thereto.

Example I
Preparation of a Genomic library

The following example illustrates the preparation of a *Bacillus* genomic library.

Genomic DNA from *Bacillus* cells is prepared as taught in Current Protocols In

Molecular Biology vol. 1, edited by Ausubel et al., John Wiley & Sons, Inc. 1987, chapter 2.

4.1. Generally, *Bacillus* cells from a saturated liquid culture are lysed and the proteins
10 removed by digestion with proteinase K. Cell wall debris, polysaccharides, and remaining
proteins are removed by selective precipitation with CTAB, and high molecular weight
genomic DNA is recovered from the resulting supernatant by isopropanol precipitation. If
exceptionally clean genomic DNA is desired, an additional step of purifying the *Bacillus*
genomic DNA on a cesium chloride gradient is added

15 After obtaining purified genomic DNA, the DNA is subjected to Sau3A digestion.
Sau3A recognizes the 4 base pair site GATC and generates fragments compatible with
several convenient phage lambda and cosmid vectors. The DNA is subjected to partial
digestion to increase the chance of obtaining random fragments.

20 The partially digested *Bacillus* genomic DNA is subjected to size fractionation on a
1% agarose gel prior to cloning into a vector. Alternatively, size fractionation on a sucrose
gradient can be used. The genomic DNA obtained from the size fractionation step is
purified away from the agarose and ligated into a cloning vector appropriate for use in a
host cell and transformed into the host cell.

Example II

The following example describes the detection of gram-positive microorganism SP1.
The same procedures can be used to detect SP2, SP3, SP4 or SP5.

25 DNA derived from a gram-positive microorganism is prepared according to the
methods disclosed in Current Protocols in Molecular Biology, Chap. 2 or 3. The nucleic acid
is subjected to hybridization and/or PCR amplification with a probe or primer derived from
SP1. A preferred probe comprises the nucleic acid section encoding conserved amino acid
residues.

30 The nucleic acid probe is labeled by combining 50 pmol of the nucleic acid and 250
mCi of [γ 32P] adenosine triphosphate (Amersham, Chicago IL) and T4
polynucleotide kinase (DuPont NEN[®], Boston MA). The labeled probe is purified with
Sephadex G-25 super fine resin column (Pharmacia). A portion containing 10^7 counts per
minute of each is used in a typical membrane based hybridization analysis of nucleic acid
sample of either genomic or cDNA origin.

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The DNA sample which has been subjected to restriction endonuclease digestion is fractionated on a 0.7 percent agarose gel and transferred to nylon membranes (Nytran Plus, Schleicher & Schuell, Durham NH). Hybridization is carried out for 16 hours at 40 degrees C. To remove nonspecific signals, blots are sequentially washed at room temperature under increasingly stringent conditions up to 0.1 x saline sodium citrate and 0.5% sodium dodecyl sulfate. The blots are exposed to film for several hours, the film developed and hybridization patterns are compared visually to detect polynucleotide homologs of *B. subtilis* SP1. The homologs are subjected to confirmatory nucleic acid sequencing. Methods for nucleic acid sequencing are well known in the art. Conventional enzymatic methods employ DNA polymerase Klenow fragment, SEQUENASE® (US Biochemical Corp, Cleveland, OH) or Taq polymerase to extend DNA chains from an oligonucleotide primer annealed to the DNA template of interest.

Various other examples and modifications of the foregoing description and examples will be apparent to a person skilled in the art after reading the disclosure without departing from the spirit and scope of the invention, and it is intended that all such examples or modifications be included within the scope of the appended claims. All publications and patents referenced herein are hereby incorporated in their entirety.

CLAIMS

1. A gram-positive microorganism having a mutation or deletion of part or all of one or more of the genes encoding a serine protease selected from the group consisting of SP1, SP2, SP3, SP4 and SP5 said mutation or deletion resulting in the inactivation of the SP1, SP2, SP3, SP4 or SP5 proteolytic activity.
2. The gram-positive microorganism according to Claim 1 that is a member of the family *Bacillus*.
3. The microorganism according to Claim 2 wherein the member is selected from the group consisting of *B. licheniformis*, *B. lentus*, *B. brevis*, *B. stearothermophilus*, *B. alkalophilus*, *B. amyloliquefaciens*, *B. coagulans*, *B. circulans*, *B. laetus* and *Bacillus thuringiensis*.
4. The microorganism of Claim 1 wherein said microorganism is capable of expressing a heterologous protein.
5. The microorganism of Claim 4 wherein said heterologous protein is selected from the group consisting of hormone, enzyme, growth factor and cytokine.
6. The microorganism of Claim 5 wherein said heterologous protein is an enzyme.
7. The microorganism of Claim 6 wherein said enzyme is selected from the group consisting of a proteases, carbohydrases, and lipases; isomerases such as racemases, epimerases, tautomerases, or mutases; transferases, kinases and phosphatases.
8. A cleaning composition comprising a serine protease selected from the group consisting of SP1, SP2, SP3, SP4 and SP5.
9. An expression vector comprising nucleic acid encoding a serine protease selected from the group consisting of SP1, SP2, SP3, SP5 and SP5.
10. A host cell comprising an expression vector according to Claim 9
11. A method for the production of a heterologous protein in a *Bacillus* host cell comprising the steps of

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(a) obtaining a *Bacillus* host cell comprising nucleic acid encoding said heterologous protein wherein said host cell contains a mutation or deletion in at least one of the genes encoding serine protease 1, serine protease 2 serine protease 3; serine protease 4 and serine protease 5.

(b) growing said *Bacillus* host cell under conditions suitable for the expression of said heterologous protein.

Rule
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12. The method of Claim 11 wherein said *Bacillus* cell is selected from the group consisting of *Bacillus subtilis*, *B. licheniformis*, *B. lentus*, *B. brevis*, *B. stearothermophilus*, *B. alkalophilus*, *B. amyloliquefaciens*, *B. coagulans*, *B. circulans*, *B. lautus* and *Bacillus thuringiensis*.

13. The method of Claim 13 wherein said *Bacillus* host cell further comprises a mutation or deletion in at least one of the genes encoding apr, npr, epr, wpr and mrp.

14. A gram-positive microorganism having at mutation or deletion in at least one of the genes encoding a serine protease selected from the group consisting of serine protease 1, serine protease 2 serine protease 3; serine protease 4 and serine protease 5.

15. The microorganism of Claim 16 further comprising a mutation or deletion in at least one of the genes encoding apr, npr, epr, wpr and mrp.

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10 30
 atggaaaagctgataaccgcagacgacatcacagcgattgtctctgt
 M K K L I T A D D I T A I V S V
 50 70 90
 accgatccctaatacgccccagacggtacccgtgccgcataatgtaaaa
 T D P Q Y A P D G T R A A Y V K
 110 130
 tcacaagtaaatcagagaaaagattcgtatacatcaaataatatggatc
 S Q V N Q E K D S Y T S N I W I
 150 170 190
 tatgaaacgaaaacgggaggatctgttccttggacacatggagaaaag
 Y E T K T G G S V P W T H G E K
 210 230
 cgaagcaccgacccaagatggtctccggacggcgacgcttgcctt
 R S T D P R W S P D G R T L A F
 250 270 290
 atttctgtatcgagaggcgatcgccacagtttatatcatgagact
 I S D R E G D A A Q L Y I M S T
 310 330
 gaaggcgaggaaagcaagaaaactgactgatatcccatatggcgtca
 E G G E A R K L T D I P Y G V S
 350 370
 aagccgctatggccccggacggtaatcgattctggtaactatcagt
 K P L W S P D G E S I L V T I S
 390 410 430
 ttgggagagggggaaagcattgtatgaccggagaaaaaacagacggagg
 L G E S I D D R E K T E Q D
 450 470
 agctatgaacctgttgaagtgcacggccttcataacacggacggc
 S Y E P V E V Q G L S Y K R D G
 490 510 530
 aaaggggctacgcgagagggtcgatggccagcttgcgttcacgcgt
 K G L T R G A Y A Q L V L V S V
 550 570
 aagtgcgggtgagatgaaagagactgacaagtccacaaagctgatcatgg
 K S G E M K E L T S H K A D H G

FIG. - 1A-1

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590 610
 gatectgtttttctctgacggcaatggcttgcgttctcagctaat
 D P A F S P D G K W L V F S A N

630 650 670
 ttaactgaaacagatgatgccagcaagccgcatgtatggcttacataatg
 L T E T D D A S K P H D V Y I M

FIG._ 1A-2

690 710
 tcactggagtcggagatctaaggcagggttacacccatcgccgctca
 S L E S G D L K Q V T P H R G S

730 750 7
 ttcggatcaagctcatttcaccagacggaaaggatcttgctttgc
 F G S S S F S P D G R Y L A L L

70 790 810
 gaaaaatgaaaaggaaatataagaatgctacgctctcaaaggcgtggc
 G N E K E Y K N A T L S K A W L

830 850
 tatgatatcgaacaaggccgcctcacatgtcttactgagatgctggac
 Y D I E Q G R L T C L T E M L D

870 890 910
 gttcatttagcggtatgcgtgatggagattcattgatcggtggct
 V H L A D A L I G D S L I G G A

930 950
 gaacacgcggccgatttggacaaaggacagccaaagggtttatgtc
 E Q R P I W T K D S Q G F Y V I

970 990 10
 ggcacagatcaaggcagttacgggcatttatatttcgattgaaggc
 G T D Q G S T G I Y Y I S I E G

10 1030 1050
 ctttgttatccgattcgctggaaaaagagtacatcaatagctttct
 L V Y P I R L E K E Y I N S F S

1070 1090
 ctttcacctgtatggacacagcactttatggccagtgacaaaggccggac
 L S P D E Q H F I A S V T K P D

FIG._ 1B-1

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1110 1130 1150
 agaccgagtgagcttacagtatcccgcttggacaggaagagaaaacag
 R P S E L Y S I P L G Q E E K Q

1170 1190
 ctgactggcgcaatgacaagttgtcagggagcatacgatatacaata
 L T G A N D K F V R E H T I S I

1210 1230 12
 cctgaagagattcaaatatgctacagaagacggcgtatggtaacggc
 P E E I Q Y A T E D G V M V N G

50 1270 1290
 tggctgtatgaggcctgcacaaatggaaaggtagacaacataccactt
 W L M R P A Q M E G E T T Y P L

1310 1330
 attcttaacatacagggcggtccgcataatgtatgtacggacatacatat
 I L N I H G G P H M M Y G H T Y

1350 1370 1390
 ttcatgagtttcagggtctggcgccgaaaggatacgcggcgtttat
 F H E F Q V L A A K G Y A V V Y

FIG._ 1B-2

1410 1430
 atcaatccgagaggaaggccacggctacgggcaggaatttgcgtaatgcg
 I N P R G S H G Y G Q E F V N A

1450 1470 14
 gtcagaggagattatggggaaaggattatgacgtatgtcaggct
 V R G D Y G G K D Y D D V M Q A

90 1510 1530
 gtggatgaggctatcaaacgagatccgcataattgatcctaagcggctc
 V D E A I K R D P H I D P K R L

1550 1570 14
 ggtgtcacggcgaaagctacggagggttatgaccaactggatcgctc
 G V T G G S Y G G F M T N W I V

1590 1610 1630
 gggcagacgaaccgccttaaagctgcgttacccagcgctcgatataca
 G Q T N R F K A A V T Q R S I S

FIG._ 1C-1

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1650 1670
 aattggatcagcttcacggcgctcagtgatatcggttatttttaca
 N W I S F H G V S D I G Y F F T

1690 1710 17
 gactggcagcttgagcatgacatgttgaggacacagaaaagctctgg
 D W Q L E H D M F E D T E K L W

30 1750 1770
 gaccgggtctccctttaaaatacgcagcaaacgtggagacaccgctttg
 D R S P L K Y A A N V E T P L L

1790 1810
 atactgcatggcgagcgggatgaccgatgcccgtcgagcaggcggag
 I L H G E R D D R C P I E Q A E

1830 1850 1870
 cagctgtttatcgctctgaaaaaaaaatgggcaaggaaaccaagcttgc
 Q L F I A L K K M G K E T K L V

1890 1910
 cgtttccgaatgcacgcacaatttatcacgcacccggacacccaaga
 R F P N A S H N L S R T G H P R

1930 1950 19
 cagcggatcaagcgctgaattatatcagctcatgggttcatcaacat
 Q R I K R L N Y I S S W F D Q H

70
 ctc
 L

FIG._1C-2

dap2_yeast	WRHSTPGSYFVYDKSSSSSEEIGNEVALAIWSPNSNDIAYQDN-NIYIYSAISKRTIRA MKKLITADDITAVSVTDPDPOYAPDGTRAAVRSQVNQEDSYTSNWIYE	170 180 190 200 210 220 10 20 30 40 50
dap2_yeast	VTNDGSSFLFNGKPDWVYEEVFEDDKAAMWSPGTYDYLAFLKIDESEVGEFFIPIPYVQDE TKTGGSV-----P-WTHGEKRSTDPR---WSPDGRTLAFISDREGDAQL---YTMSTE	230 240 250 260 270 280 60 70 80 90
dap2_yeast	KDIYFEMRSIKYKPSKG---TPNPHAEILWYSMKGDTSFHPRISGNKKG---SLLITEVTVW GGEARKLTDIPIGVSKPLWSPDGESILVITISIGEGESIDDR-EKTEQDSYEPVEVQGLSY	290 300 310 320 330 330 100 110 120 130 140 150
dap2_yeast	VGNGNVLYVKTDRSSDLITVFLDITIAKTSNVRNE----SNGGGWWEITHNTLFIPANE KRDGKGGLTRGAYAQOLVLSVKSGEMKELTSKADHGDPAFSPDGKWLVESAN---LTETD	340 350 360 370 380 390 160 170 180 190 200 210
dap2_yeast	TFDRPHNGYYDILPIGGYN----HLAYFENSNS---HYKTLTEGKWEVNGPLA---F DASEPHDVYIMSLSESGLIKQVTPHRGSFGSSSSPDPGRYLALIGNEKEYKNATLSKAWLY	400 410 420 430 440 440 220 230 240 250 260 270
dap2_yeast	DSMENRLYFISTRKSSTERHYYID-LRSPNIEIYVTDSEGDGVYDVSTSSGRRFLG---L DIEGRLTCLEMLDVHADLIGDSLIGGAEQRPIWTKDSQ96FVIGTDQGST-GIYI	450 460 470 480 490 499 280 290 300 310 320 330

FIG.-2A

dap2_yeast	500	510	520	530	540	550
	TYKGPKVQYQKIVDFHSRKAERCDKGNVIGKSLYHLEKNEVLTTRILEDYAVPR-KSFREL					
YUXL	: :	: :	: :	: :	: :	
	SIEGLVYPIRLEKEYTINSFSLSDEQHFTASVTKPDRSEL-----YSIPLGQEERKQL					
	340	350	360	370	380	
dap2_yeast	560		570	580	590	600
	NLGKDFEGKD-----ILVNSEYLIPNDFEDTLSHYPPVFFFAYGGPNSQ					
YUXL	: : :	: : :	: : :	: : :	: : :	
	TGANDKFVREHTISLPEEIQYATEDGVMVNGMLMRPAQMEGETT--YPLLNTHGGPH-M					
	390	400	410	420	430	440
dap2_yeast	610	620	630	640	650	660
	QVVRTEFSGVNFNEVVAQSLQNLNAIVVUDGRTGFKQDFRSVLVRDLRGDYEARDQIISAS-L					
YUXL	: :	: : :	: : :	: : :	: : :	
	MYGHTYFHEEF-QVLAARCYA-VVYINPRSHGXYGOEVNAVRGDYGGDYDDVMQAVDEA					
	450	460	470	480	490	
	↓ Ser					
dap2_yeast	670		680	690	700	710
	YGSLTFVDPKQKISLFGNSYGGYLTKLTKDGGGRFKEYGMSVAPVTDRFYDSVTERYM					
YUXL	: : :	: : :	: : :	: : :	: : :	
	IKRDPHIDPRLGVTGGSYGGFMNTNWIGOTN--RFKAVIATRSISNWSFHGVSIDGYF					
	510	520	530	540	550	
dap2_yeast	730		740	750	760	770
	HTP-QENFDGYVES-SVHNVTALAQNRR---FLIMHGTGDDNVHFQNSLKFEDLDLNG					
YUXL	:	: :	: :	: :	: :	
	FTDWOLEHDMEFDTEKLWDRSPSLPKYAANVETPLLHGERDRCPIQEAEQLFIAALKKMG					
	560	570	580	590	600	610
	↓ His					
dap2_yeast	780	790	800	810		
	VENYDVHVFPPDSDHSIRYHANANIVFDKLIDWKAFAFDGQFVK					
YUXL	:	:	:	:		
	KETKLVR-FPMASHNLNSRTGHPPQRIKRNYISSLWDQHL					
	620	630	640	650		

FIG..2B

380	390	400	410	420	430	439
yux1.bsupep	QEEKQLTGANDKFVRREHTISIPEEIOYATEDGVMVNQWLMRPAQMGETTYPLILNTHGG					
YTM1	MIVEKRRFPSPSQHVRILYTCIYLSNLRVKGLLAEPAE-PGQ--YDGFLYLRGG					
	10	20	30	40	50	
440	450	460	470	480	490	
yux1.bsupep	PHMMYGHYTFHEQVLAKGAYAVVYINPRGSHG-YGQEFVNARVGDYGGDDVMQAVD					
YTM1	IKSV-GMVRPGRIIQFASQEFVVFAPEYRGNGQEGNE-----DFAGEDREDAFSAF-					
	60	70	80	90	100	
500	510	520	530	540	550	
yux1.bsupep	EAIKRDPHIDPKRLGTVGTYGGEMTNWIVGOTNREFFKAUTORSISNWISFHGVSDIGYF					
YTM1	RLQQHPNVKKDRIHIFGSRGGM-----GMLTAIENGQQAASEVSW--GGVSDMILT					
	110	120	130	140	150	
560	570		580	590	600	
yux1.bsupep	FTDWQLEHDMFEDT-----EKLWDRSPPLKYAANVETPLLIHGERRDRCPIEQAE					
YTM1	YEERQDILRMMKRVIGGTPKKVPEEYQW-RTPFDQVNKIQAPVLLHGERDQNVSIQHSY					
	160	170	180	190	200	
610	620	630	640	650		
yux1.bsupep	QLFTIALKKMGKRETKLVRFFNASHNLSRTGHPRQRIRKINYISSWFQHL					
YTM1	LIEEKIKQLHKRVETWYISTFTHP---PKENRVRVRLTOMKRN					
	220	230	240	250		

FIG.-3

yux1.bsupep	PEEIQAYATEDGVMVNGWLMRPAQMEGETTYPLILINHGGPHMMYGHTYFHEFOVLAARKGY	410	420	430	440	450	460
Y1TV	MIQIENQTVSGIPFLHVKEENRHRAPLVYFIHGFTSAKE-HN-LHIAVILLAERKGF	10	20	30	40	50	
yux1.bsupep	AVVYINPRSHGYGQEFVNAVRGDGYKDDYDMMQANDEA-----IKRDPHIDDPKRQLGV	470	480	490	500	510	
Y1TV	RAVL--PEALH-HGERGEMAAVEELAGHFWDIVLNEEEEIGVULKNHFEKEGLIDGGRIGL	60	70	80	90	100	110
yux1.bsupep	TGGSYGGMNTNWIGTQNTNRKFAAVTORSISNWISFHGSIDGYFTTWOLEHMFD-TE	520	530	540	550	560	570
Y1TV	AGTSMGGITTLGALTAYDWIKAGVSLMSPNVYELFQ-QQIDHI-QSQGIEIDVPEEKVQ	120	130	140	150	160	170
	Ser↑						
yux1.bsupep	KLWDRSPLKYYAANV----EPLPLILHGGERDDRCPIEQAEQOLFIAKKMGKET---KLV	580	590	600	610	620	
Y1TV	QLMKRLEFLDSLQPEKQQLQRPPLFWHGAKDKVVPYAPTEKFYDTIKSHYSEQPERLQFI	180	190	200	210	220	230
	Asp↑						
yux1.bsupep	RFPNASHNLSRTGHPRQRIRKRNIVSSMFQDQLL	630	640	650	660	670	
Y1TV	GDENAADHKV-----PRAAV--LKTIE-WFETYL	240	250				
	↑Hs						

FIG.-4

yux1.bsupep	390	400	410	420	430	440
YQKD	TIKRETNDGHDVESFEQMEKTAFVIPSAYGDIKGYHIVAPHDTPTNTLICIGVTMVNLN	40	50	60	70	80
yux1.bsupep	450	460	470	480	490	500
YQKD	GHTYFHEFOVLAARCYAVVYINPRGSHGGQEFVNARVGDYGGKDYDDVMQAVDEAIKRD	100	110	120	130	140
yux1.bsupep	510	520	530	540	550	559
YQKD	PHIDPKRLGVTGGSYGGFMNIVGQ-----TNRFEKAUTQRSISNWISFHGVSDIGYFF	150	160	170	180	190
yux1.bsupep	560	570	580	590	600	610
YQKD	NHRG--LIGHGESMGAVTALLYAGAHCSGDAFYIADCPFACFDEQLAYLRAE--YRL	210	220	230	240	250
yux1.bsupep	620	630	640	650		
YQKD	KKRGPKALYIA-ENGEHANSYTKNRHTYRKTQEFLLDDNNNDSTE	270	280	290	300	

↑ His **FIG.-5**

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10 30
ttgatttagagaaaaagaagattccgtcgccaaaggcagcatgtgcgt
L I V E K R R F P S P S Q H V R

50 70 90
ttgtatacgatctgtatctgtcaaatggattacgggtaaggggctt
L Y T I C Y L S N G L R V K G L

110 130
ctggctgagccggcggaaaccggacaatatgacggattttatatttg
L A E P A E P G Q Y D G F L Y L

150 170 190
cgcggcgggattaaaagcgtggcatggttcggccggccggattatc
R G G I K S V G M V R P G R I I

210 230
cagtttgcattccaaagggtttgtgtgtttgtctttttacagaggc
Q F A S Q G F V V F A P F Y R G

250 270 2
aatcaaggaggagaaggcaatgaggatttgcggagaagacaggag
N Q G G E G N E D F A G E D R E

90 310 330
gatgcatttctgtttcgcttcgttcagcagcacccaaatgtcaag
D A F S A F R L L Q Q H P N V K

350 370
aaggatagaatccatatcttcggttttccgcgggaattatggga
K D R I H I F G F S R G G I M G

390 410 430
atgctactgcgtcgaaatggcggcaggcagcttcattgtttcc
M L T A I E M G G Q A A S F V S

450 470
tggggaggcgtcagtgtatgattttcacatcggaggcggcaggat
W G G V S D M I L T Y E E R Q D

490 510 5
ttgcggcgaatgtatgaaaaagagtcatcgccggaaacaccgaaaaaggtg
L R R M M K R V I G G T P K K V

30 550 570
cctgaggaatatcaatggaggacaccgtttgaccaactaaacaaaatt
P E E Y Q W R T P F D Q V N K I

FIG.-7A

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590 610
caggctcccgctgtttaatccatggagaaaaagacccaaaatgttcg
Q A P V L L I H G E K D Q N V S

630 650 670
attcagcattcctatttattagaagagaagctaaaaacaactgcataag
I Q H S Y L L E E K L K Q L H K

690 710
ccgggtggaaacatggtactacatgtacattcacacattttccgc
P V E T W Y Y S T F T H Y F P P

730 750 7
aaagaaaaaccggcgatcgatcgatcgccagctcacacaatggatgaaaaac
K E N R R I V R Q L T Q W M K N

70
cgc
R

FIG._7B

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10 30
 gtgatacaaattgagaatcaaaccgttccggatattccgttttacat
 V I Q I E N Q T V S G I P F L H

50 70 90
 attgtaaaggaagagaacaggcaccgcgcgtgtccctctcgatctt
 I V K E E N R H R A V P L V I F

110 130
 atacatggtttacaaggcgcgaaggaaacacaacccatattgcttat
 I H G F T S A K E H N L H I A Y

150 170 190
 ctgcttcggagaagggttttagagccgttctgcggaggcttgcac
 L L A E K G F R A V L P E A L H

210 230
 catggggaaacggggagaagaaaatggctgttgaagagagctggcgccat
 H G E R G E E M A V E E L A G H

250 270 2
 ttttggatatcgcccaacagagattgaagagatcggctacttaaa
 F W D I V L N E I E E I G V L K

90 310 330
 aaccattttggaaaaagagggcctgatagacggcgccgcattggcttc
 N H F E K E G L I D G G R I G L

350 370
 gcaggcacgtcaatggcgccatcacacgcgttggcttgcata
 A G T S M G G I T T L G A L T A

390 410 430
 tatgattggataaaagccggcgtcaggctgatggaaagccgattac
 Y D W I K A G V S L M G S P N Y

450 470
 gtggagctgtttcagcagcaggattgaccatattcaatctcaggcatt
 V E L F Q Q I D H I Q S Q G I

490 510 5
 gaaaatcgatgtccggaaagagaaggtaacagcagctgatgaaacgttc
 E I D V P E E K V Q Q L M K R L

30 550 570
 gagttgcggatctcaggcttcagccggagaaaactgcaacagcggcc
 E L R D L S L Q P E K L Q Q R P

FIG._8A

590

cttttatttggcacggcgcaaaagataaaagtgtgccttacgcgccc
L L F W H G A K D K V V P Y A P

630

650

670

acccggaaattttatgacacgattaaatcccattacagcgagcagccg
T R K F Y D T I K S H Y S E Q P

690

gaacgcctgcaatttatcgagataaaaacgctgaccataaagtcccg
E R L Q F I G D E N A D H K V P

730

750

cgggcagctgttaaaaacgattgaatggtttgaaacgtactta
R A A V L K T I E W F E T Y L

FIG._8B

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10 30
ttgaagaaaaatcctttggccattggcgcgctcgtaacagctgtcatc
L K K I L L A I G A L V T A V I

50 70 90
gcaatcggaatttttcacatcatgattctattcatcaagaaaaaa
A I G I V F S H M I L F I K K K

110 130
acggatgaagacattatcaaaagagagacagacaacggacatgtg
T D E D I I K R E T D N G H D V

150 170 190
tttgaatcatttgaacaaatggagaaaaccgctttgtgataccctcc
F E S F E Q M E K T A F V I P S

210 230
gcttacgggtacgacataaaaggataccatgtcgcacccgcatgacaca
A Y G Y D I K G Y H V A P H D T

250 270 2
ccaaataccatcatcatctgccacgggtgacgatgaatgtactgaat
P N T I I I C H G V T M N V L N

90 310 330
tctcttaagtatatgcatttattctagatctcggtggaaatgtgctc
S L K Y M H L F L D L G W N V L

350 370
atttatgacccatcgccggatggccaaagcgccggaaagacgaccagc
I Y D H R R H G Q S G G K T T S

390 410 430
tacgggttttacgaaaaggatgatctaataaggttgcagctgtctc
Y G F Y E K D D L N K V V S L L

450 470
aaaaacaaaaacaaatcatcgccgattgtatcgaaattcatggtgagtcg
K N K T N H R G L I G I H G E S

490 510 5
atggggccgtgaccgcctgtttatgtctggtgacactgcagcgt
M G A V T A L L Y A G A H C S D

30 550 570
ggcgctgattttatattgcgcattgtccgtcgatgtttgtgaa
G A D F Y I A D C P F A C F D E

FIG._9A

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590 610
 cagcttgcctatcggtctgagagcggaaatacaggctccgtcttggccc
 Q L A Y R L R A E Y R L P S W P

630 650 670
 ctgcttcctatcgccgacttcttttgaagctgaggggaggctatcg
 L L P I A D F F L K L R G G Y R

690 710
 gcacgtgaagtatctccgttgcattgtcattgataaaaattgaaaagccg
 A R E V S P L A V I D K I E K P

730 750 7
 gtccttttattcacagtaaggatgtactacattcctgtttctca
 V L F I H S K D D D Y I P V S S

70 790 810
 acccgagcggctttatgaaaagaaaacgcggccgaaagcgctgtacatt
 T E R L Y E K K R G P K A L Y I

830 850
 gccgagaacggtaaacacgcgtatgtcatataccaaaaatcgccatacg
 A E N G E H A M S Y T K N R H T

870 890 910
 taccgaaaaacagtgcaggagtttttagacaacatgaatgattcaaca
 Y R K T V Q E F L D N M N D S T

gaa
 E

FIG._9B

DECLARATION
AND POWER OF ATTORNEY

AS A BELOW NAMED INVENTOR, I HEREBY DECLARE THAT:

MY RESIDENCE, POST OFFICE ADDRESS AND CITIZENSHIP ARE AS STATED BELOW NEXT TO MY NAME. I BELIEVE I AM THE ORIGINAL, FIRST AND SOLE INVENTOR (IF ONLY ONE NAME IS LISTED BELOW) OR AN ORIGINAL, FIRST AND JOINT INVENTOR (IF PLURAL NAMES ARE LISTED BELOW) OF THE SUBJECT MATTER WHICH IS CLAIMED AND FOR WHICH A PATENT IS SOUGHT ON THE INVENTION ENTITLED **PROTEASES FROM GRAM-POSITIVE ORGANISMS** THE SPECIFICATION OF WHICH

CHECK ONE:

IS ATTACHED HERETO
WAS FILED ON _____ AS APPLICATION SERIAL NO. _____ AND WAS AMENDED ON _____.

I HEREBY STATE THAT I HAVE REVIEWED AND UNDERSTAND THE CONTENTS OF THE ABOVE IDENTIFIED SPECIFICATION, INCLUDING THE CLAIMS, AS AMENDED BY ANY AMENDMENT REFERRED TO ABOVE. I ACKNOWLEDGE THE DUTY TO DISCLOSE INFORMATION WHICH IS MATERIAL TO PATENTABILITY AS DEFINED IN TITLE 37, CODE OF FEDERAL REGULATIONS §1.56.

I HEREBY CLAIM FOREIGN PRIORITY BENEFITS UNDER TITLE 35, UNITED STATES CODE §119, OF ANY FOREIGN APPLICATION(S) FOR PATENT OR INVENTOR'S CERTIFICATE LISTED BELOW AND HAVE ALSO IDENTIFIED BELOW ANY FOREIGN APPLICATION FOR PATENT OR INVENTOR'S CERTIFICATE HAVING A FILING DATE BEFORE THAT OF THE APPLICATION ON WHICH PRIORITY IS CLAIMED.

APPLICATION NUMBER	COUNTRY	DATE OF FILING	PRIORITY CLAIMED YES NO

I HEREBY CLAIM THE BENEFIT UNDER TITLE 35, UNITED STATES CODE §120, OF ANY UNITED STATES APPLICATION(S) OR PCT INTERNATIONAL APPLICATION(S) DESIGNATING THE UNITED STATES OF AMERICA THAT IS LISTED BELOW AND, INSOFAR AS THE SUBJECT MATTER OF EACH OF THE CLAIMS OF THIS APPLICATION IS NOT DISCLOSED IN THE PRIOR UNITED STATES APPLICATION IN THE MANNER PROVIDED BY THE FIRST PARAGRAPH OF TITLE 35, UNITED STATES CODE §112, I ACKNOWLEDGE THE DUTY TO DISCLOSE MATERIAL INFORMATION AS DEFINED IN TITLE 37, CODE OF FEDERAL REGULATIONS §1.56(a) WHICH OCCURRED BETWEEN THE FILING DATE OF THE PRIOR APPLICATION AND THE NATIONAL OR PCT INTERNATIONAL FILING DATE OF THIS APPLICATION.

APPLICATION NUMBER	DATE OF FILING	STATUS - PATENTED, PENDING OR ABANDONED
PCT/US98/14647	14 JULY 1998	PENDING
EP 97305232.7	15 JULY 1997	ABANDONED

POWER OF ATTORNEY: AS A NAMED INVENTOR I HEREBY APPOINT AS MY ATTORNEY(S) WITH FULL POWER OF SUBSTITUTION AND REVOCATION, TO PROSECUTE THIS APPLICATION AND TRANACT ALL BUSINESS IN THE PATENT AND TRADEMARK OFFICE CONNECTED THEREWITH:

(L)

MARGARETA A. HORN, REG. NO. 33,401;
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FULL NAME OF INVENTOR	FULL FIRST NAME DAVID	INITIAL A	LAST NAME ESTELL
RESIDENCE & CITIZENSHIP	CITY SAN MATEO,	STATE OR FOREIGN COUNTRY CALIFORNIA	COUNTRY OF CITIZENSHIP USA
POST OFFICE ADDRESS	POST OFFICE ADDRESS 248 WOODBRIDGE CIRCLE,	CITY SAN MATEO	STATE OR COUNTRY CALIFORNIA
			ZIP CODE 94403

I FURTHER DECLARE THAT ALL STATEMENTS MADE HEREIN OF MY OWN KNOWLEDGE ARE TRUE AND THAT ALL STATEMENTS MADE ON INFORMATION AND BELIEF ARE BELIEVED TO BE TRUE; AND FURTHER THAT THESE STATEMENTS WERE MADE WITH THE KNOWLEDGE THAT WILLFUL FALSE STATEMENTS AND THE LIKE SO MADE ARE PUNISHABLE BY FINE OR IMPRISONMENT, OR BOTH, UNDER SECTION 1001 OF TITLE 18 OF THE UNITED STATES CODE, AND THAT SUCH WILLFUL FALSE STATEMENTS MAY JEOPARDIZE THE VALIDITY OF THE APPLICATION OR ANY PATENT ISSUING THEREON.

SIGNATURE OF INVENTOR 201	
DATE	January 5, 2000